WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)			
(51) International Patent Classification 7:		(11) International Publication Number: WO 00/10526	
A61K 9/00	A2	(43) International Publication Date: 2 March 2000 (02.03.00)	
(21) International Application Number: PCT/	EP99/060	Pommard, F-75012 Paris (FR). ZÜGER, Othmar [CH/CH]; Heuwinkelstrasse 13, CH-4123 Allschwil (CH).	
(22) International Filing Date: 19 August 1999	9 (19.08.9	(74) Agent: BECKER, Konrad; Novartis AG, Corporate Intellectual Property, Patent & Trademark Dept., CH-4002 Basel (CH).	
(30) Priority Data: 9818340.3 9823477.6 9810320.2 9810320.2 9811059.5 (71) Applicant (for all designated States except AT US) TIS AG [CH/CH]; Schwarzwaldallee 215, CH-(CH). (71) Applicant (for AT only): NOVARTIS-ERFINDUN WALTUNGSGESELLSCHAFT MBH [AT/AT Strasse 59, A-1230 Vienna (AT).	98) C C C P: NOVAI -4058 Bas	YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE),	
 (72) Inventors; and (75) Inventors/Applicants (for US only): DE BRU [NL/FR]; 13, rue du Muhlberg, F-68730 (FR). ENGEL, Günter [DE/DE]; Im Haser D-79576 Weil (DE). PFANNKUCHE, F [DE/DE]; Vierthauen 17, D-79576 Weil (DE) SEN, Michael [DE/DE]; Apollinarisstrasse 26 	Blotzheingarten l Hans-Jürge THEWI	m Without international search report and to be republished upon receipt of that report. S-	

Published

(54) Title: NEW ORAL FORMULATION FOR 5-HT4 AGONISTS OR ANTAGONISTS

SEN, Michael [DE/DE]; Apollinarisstrasse 26, D-40227 Düsseldorf (DE). VITZLING, Christian [FR/FR]; 39, rue de

The present invention relates to a pharmaceutical composition, in particular to a composition for administering active agents which are poorly soluble in aqueous media, and/or which are acid sensitive.

> U.S. Serial No.: 09/900,336 Docket No.: 484482000300

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Słovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	T)	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

NEW ORAL FORMULATION FOR 5-HT4 AGONISTS OR ANTAGONISTS

The present invention relates to a pharmaceutical composition, in particular to a composition for administering active agents which are poorly soluble in aqueous media and/or which are acid sensitive. More particularly, the present invention relates to a pharmaceutical composition for administering active agents acting on the gastro-intestinal system. The present invention also relates to a process for manufacturing such compositions. The term "pharmaceutical" also covers veterinary use.

Pharmaceutical compositions containing active agents which are poorly soluble in aqueous media and/or acid sensitive are difficult to manufacture. One of the problems that may occur concerns adsorption of the active agent on the process equipment during the manufacturing process. Due to the poor solubility of such active agents it is also difficult to obtain pharmaceutical compositions which upon administration have a good dissolution rate. As a further problem, active agents may be degraded, e.g., chemically, during a manufacturing process using acidic conditions or during the storage of the composition.

The present invention provides compositions and processes which avoids or minimise one or more of the above problems.

We have now surprisingly found that it is possible to produce a pharmaceutical composition for administering of active agents which are poorly soluble in aqueous media, e.g., pure water, and/or acid sensitive, and which upon administration has good dissolution properties, a good bioavailability and is surprisingly efficacious.

The present invention provides in one aspect a solid oral pharmaceutical composition, e.g., a tablet, comprising an active agent which is poorly soluble in aqueous media, and/or acid sensitive, and a disintegrant, e.g., a super-disintegrant, which is present in an amount of at least 15% by weight based on the total weight of the composition.

By "poorly soluble" is meant an active agent having a solubility in aqueous media more than 0.001% and less than 10%, e.g., less than 1 %, .g., less than 0.1%, e.g., less than 0.05%, e.g., less than 0.02%, at room temperature, e.g., 25°C.

By "acid sensitive" is meant an active agent which under even slightly acidic conditions, e.g., at pH 6, may be transformed to a significant extent in a degradation product, e.g., by chemical degradation, which may have no or changed activity, e.g., within 2 hours. Examples of compounds are known in the art and may be ascertained by routine experimentation.

By "disintegrant" is meant a substance or mixture of substance added to a solid pharmaceutical composition, e.g., a tablet, to facilitate its break-up or disintegration after administration in order that the active ingredient is released from the composition as efficiently as possible to allow for its rapid dissolution (see e.g. "Remington's Pharmaceutical Science" 18th edition (1990), "The Theory and Practice of Industrial Pharmacy" Lachman et al. Lea & Febiger (1970)).

We have also found difficulties on producing stable commercially acceptable formulations, e.g., tablets, of compounds such as those disclosed in EP505322 (herein incorporated by reference) and which are useful as 5-HT₄ receptor agonists or partial agonists.

A preferred 5-HT₄ partial agonist disclosed in EP505322 is Tegaserod (3-(5-methoxy-1H-indol-3-yl-methylene)-N-pentylcarbazimidamide) (example 13) of formula

$$CH_3$$

$$CH = N - N - C - NHC_5H_{11}$$

$$NH$$

which is referred hereinafter as Compound A, or a pharmaceutically acceptable salt form thereof, e.g., the hydrogen maleate (hereinafter "hml") salt. Compound A has a solubility of about 0.02% at 25°C in water and is acid sensitive. We have found that compositions may be produced which give good absorption even in the stomach. We have also found that Compound A may be adsorbed by certain excipients so that its dissolution upon administration may be substantially reduced.

Little has been published in detail on 5-HT₄ receptor agonists, partial agonists or antagonists biopharmaceutical properties, e.g., their site of action is not known.

The present invention provides in a further aspect pharmaceutical compositions allowing a complete dissolution of 5-HT₄ receptor agonists, partial agonists or antagonists, e.g., Compound A, when administered to humans, e.g., patients, in need thereof. These compositions allow a good bioavailability and are surprisingly efficacious. Moreover, they are stable and well reproducible. A process for their preparation is also provided.

Active agents which may be used in compositions according to the present invention are more generally those acting on the gastro-intestinal system, e.g., serotonergic active agents, e.g., full agonists, partial agonists and antagonists of 5-HT₄ receptors to the extent they are poorly soluble and/or acid sensitive. They are preferably in salt form, e.g., hydrogen maleate or hydrochloride, and may be in free form.

The 5-HT₄ receptor is a cloned species of the serotonin receptor family which comprises at least 14 distinct G protein-coupled receptors (the receptor ionophore of the 5-HT₃ subtype excluded). Four splice variants of the human receptor, 5-HT_{4A}, 5-HT_{4B}, 5-HT_{4C}, and 5-HT_{4D}, have been identified which differ in the length and sequence of the protein's C terminus (Blondel et al., FEBS Letters (1997) 412:465-474; Blondel et al., J. Neurochem. (1998) 70: 2252-2261). Biochemical characterisation of 5-HT₄ receptors revealed a positive coupling to adenylyl cyclase. 5-HT₄ receptor expression in man has been found in the brain, the gut, the atria, the urinary bladder and kidneys.

Compounds capable of acting on the serotonin receptor are substituted benzamides, *e.g.*, cisapride, renzapride, zacopride, clebopride, cinitapride, mosapride, lintopride, metoclopramide, or benzoic esters, *e.g.*, RS 23597-190, SB 204070, SB 207710, or aminoguanidines, zacopride, prucalopride, SB 205149, SC 53116, RS 67333, RS 67506, BIMU 1, BIMU 8, (S)-RS 56532, Tropisetron, Alosetron, GR 113808, GR 125487, SB 207266, RS 23597, RS 39604, RS 100235, DAU 6285, SC 53606, 3-(5-hydroxy-7-methyl-1H-indol-3-yl-methylene)-N-pentyl-N-methyl-carbazimidamide, indazole-3-carboxamides, 2-oxobenzamidazole-3-carboxamides (as disclosed in EP 908 459 which is herein incorporated by referenc) etc.

5-HT₄ receptor agonists are considered as compounds which can activate 5-HT₄ receptors under quiescent/resting conditions (complete or partial activation). As 5-HT₄ receptor full agonists or partial agonists one may cite (S)-zacopride, cisapride, prucalopride, SB 205149, SC 53116, RS 67333, RS 67506, BIMU 1, BIMU 8, (S)-RS 56532 and Compound A, particularly its hydrogen maleate salt.

5-HT₄ receptor antagonists are considered as compounds which do not activate 5-HT₄ receptors but act as inhibitors of agonists at 5-HT₄ receptors. As 5-HT₄ receptor antagonists one may cite GR 113808, GR 125487, SB 203186, SB 204070, SB 207266, RS 23597, RS 39604, RS 100235, DAU 6285, SC 53606, 3-(5-hydroxy-7-methyl-1H-indol-3-yl-methylene)-N-pentyl-N-methyl-carbazimidamide.

5-HT₄ receptor agonists are useful for the prevention and treatment of gastro-intestinal motility disorders, e.g., Irritable Bowel Syndrome (IBS), Gastro-Esophageal Reflux Disease (GERD), Functional Dyspepsia (FD) and Post Operative Ileus (POI).

In a preferred embodiment, the composition of the invention comprises 20 to 60%, e.g., 30 to 50%, e.g. 40% by weight of disintegrant based on the total weight of the composition. We have observed that the use of such a high percentage of disintegrant further improves the dissolution rate in aqueous media, but also prevents the active agent from adsorbing on excipients.

As disintegrants the composition of the invention may comprise:

- crospovidone (molecular weight >10⁶), e.g., Polyplasdone XL[®], Kollidon CL[®], Polyplasdone XL-10[®].
- pregelatinised starch (MW: 30 000 120 000), e.g., starch 1500°, STA-Rx 1500°,
- sodium starch glycolate (MW: 500 000 1 000 000), e.g. Primojel®,
- carboxymethylcellulose calcium (CMC-Ca),
- carboxymethylcellulose sodium (CMC-Na) (MW: 90 000 700 000), e.g., Ac-Di-Sol®,
- sodium alginate,
- or a mixture thereof.

Preferably, the disintegrant is crospovidone which is preferably water insoluble. Preferably it rapidly exhibits high capillary or pronounced hydration capacity with little tendency to gel formation. Preferably the particle size is from about 1 to 500 micrometers. Preferred particle size distribution is less than 400 micrometers, e.g., for Polyplasdone XL®, less than 80 micrometers, e.g., less than 74 micrometers for, e.g., Polyplasdone XL-10®, approximately 50% greater than 50 micrometers and maximum of 1% greater than 250 micrometers in size for, e.g., Kollidon CL®. A preferred crospovidone is Polyplasdone XL®, e.g., with a density of about 0.213 g/cm³ (bulk) or 0.273 g/cm³ (tapped).

The pharmaceutical composition of the invention may further comprise one or more excipients.

The composition may further comprise one or more lubricants, e.g., in an amount within the range of from, e.g., 1 to 20%, e.g., from 5 to 15%, e.g., 10% by weight of the composition.

Examples of such lubricants include

- glyceryl mono fatty acid, e.g., having a molecular weight of from 200 to 800, e.g., glyceryl monostearate (e.g., Myvaplex®, USP quality)
- polyethylene glycol (PEG), having a molecular weight of from 100 to 10000, e.g., 1000 to 8000, e.g., 2000 to 6000, e.g., 2500 to 5000, e.g., Macrogol 4000 (Pulver) BP,
- hydrogenated castor oil (e.g., Cutina®), and the like or a mixture thereof.

In a preferred composition the lubricant is glyceryl monostearate. The lubricant properties of such preferred composition may be improved by adding polyethylene glycol (PEG), e.g., Macrogol 4000 (Pulver) BP.

The composition of the invention may comprise one or more surfactants, e.g., in an amount in the range of from 0.1 to 10%, e.g., 1 to 5%, e.g. 2% by weight of the total composition. Pharmaceutically suitable surfactants may be non-ionic or anionic.

As non-ionic surfactants one may use:

- polyoxyethylene-sorbitan-fatty acid esters (polysorbates; MW: 500 to 2000), e.g., mono-and tri- lauryl, palmityl, stearyl and oleyl esters, e.g., Tween®, e.g., Tween 80®;

- polyoxyethylene fatty acid esters (MW: 500 to 5000), e.g., Myrj® or Cetiof®;
- polyoxyethylene-polyoxypropylene co-polymers, e.g., having a molecular weight of from 1000 to 20 000, e.g., 6 000 to 15 000, e.g., 7 000 to 10 000, e.g., Pluronic® or Emkalyx®;
- polyoxyethylene-polyoxypropylene block co-polymers e.g., having a molecular weight of from 1000 to 20 000, e.g., 6 000 to 15 000, e.g., 7 000 to 10 000, e.g., Poloxamer 188[®];
- reaction products of a natural or hydrogenated castor oil and ethylene oxide, e.g., Cremophor®;
- dioctylsuccinate or di-[2-ethylhexyl]- succinate;
- propyleneglycol mono- and di-fatty acid (e.g. C₆-C₈) esters, e.g., Miglyol[®]; or mixtures thereof.

As suitable anionic surfactants one may use, e.g., sodium laurylsulfate or docusate sodium.

Unless where otherwise stated fatty acid or carbon containing chain is from about 8 to 22 carbon atoms, e.g., C_{18} .

The composition of the invention may comprise one or more binders, e.g., in an amount in the range of from 1 to 10%, e.g., 2 to 8%, e.g., 5% by weight. One may particularly use:

- hydroxypropylmethylcellulose, e.g., having a molecular weight of from 10 000 to 1 500 000, e.g., HPMC-3 (3mPa-s) (e.g. Pharmacoat®, Methocel®),
- polyvinylpyrrolidone, e.g., having a molecular weight of from 2500 to 3 000 000, e.g., 8 000 to 1 000 000, e.g., 10 000 to 400 000, e.g., 30 000 to 50 000 (e.g., Kollidon $^{\$}$, Plasdone $^{\$}$),
- potato starch, wheat starch, com starch, e.g., having a molecular weight of from 30 000 to 120 000,

or a mixture thereof.

The composition of the invention may comprise one or more diluents such as lactose, mannitol, sucrose, calcium sulphate, calcium phosphate, microcristalline cellulose (Avicel®) in an amount within the range of from, e.g., 10 to 70%, e.g., 20 to 50%, e.g., 30% by weight of the composition. Preferably, the diluent is lactose, e.g., lactose 200 mesh (e.g., from DMV® or Alpavit ®), e.g., the monohydrated form.

Other conventional excipients which may optionally be present in the composition of the invention include preservatives, stabilisers, anti-adherents or silica flow conditioners or glidants, e.g., silicon dioxide (e.g., Syloid[®], Aerosil[®]) as well as FD&C colours such as ferric oxides.

Other excipients disclosed in the literature, as for instance in Fiedler's "Lexicon der Hilfstoffe", 4th Edition, ECV Aulendorf 1996 and "Handbook of Pharmaceutical Excipients" Wade and Weller Ed.(1994), the contents of which are incorporated herein by reference, may be used in the pharmaceutical compositions according to the invention.

The invention is particularly useful for pharmaceutical compositions containing an active agent, e.g., an 5HT₄ receptor agonist, partial agonist or antagonist, e.g., compound A, e.g., the hydrogen maleate salt, which is present in an amount within the range of from about 0.2% to about 20%, e.g. 0.5 to 15%, and preferably from about 1% to about 10% by weight of the composition.

A preferred composition of the invention may comprise from about 0.5 to about 15% by weight of active agent, e.g., a 5HT₄ receptor agonist, e.g., compound A, e.g., the hydrogen maleate salt, from 20 to 60% by weight of disintegrant, e.g., crospovidone, from 1 to about 20% by weight of a lubricant, e.g., monoglycerylstearate, from 0.1 to about 10% by weight of a surfactant, e.g., poloxalkol, from about 10 to 50% by weight of a diluent, e.g., lactose, and from 1 to 10% by weight of a binder, e.g., hydroxypropylmethyl cellulose (e.g. HPMC-3). From 1 to 10% by weight of PEG may also be added.

The weight ratio of the active agent to the disintegrant may be from 1:1 to 1:400, e.g., 1:5 to 1:100, 1:8 to 1:50, e.g., 1:16 to 1:20.

In a further aspect the present invention provides a pharmaceutical oral, e.g., tablet, composition comprising one of the active agents cited above, e.g., a 5-HT₄ agonist, partial agonist or antagonist, e.g., Tegaserod, said composition having dissolution characteristics in water or in USP buffers pH 6.8 and 7.5 of:

time (minutes)

amount (percentage)

5

15	80 - 100
30	95 - 100
60	100

For example a composition according to the invention, e.g., comprising Tegaserod as the active agent, may have dissolution characteristics in water or in USP buffers pH 6.8 and 7.5 of:

time (minutes)	amount (percentage)
5	48.9
15	95.5
30	99.7
60	100

In a further aspect the present invention provides a pharmaceutical oral, e.g., tablet, composition comprising one of the active agents cited above, e.g., a 5-HT₄ agonist, partial agonist or antagonist, e.g., Tegaserod, wherein in use 80% of said active agent is released in water or in USP buffers pH 6.8 and 7.5 within 5 minutes.

In a further aspect, the present invention provides the use of at least 15% by weight of a disintegrant in the manufacturing of pharmaceutical composition for the administration of an acid sensitive and/or poorly soluble, e.g., in aqueous media, active agent, e.g., a 5-HT₄ receptor agonist, e.g., compound A, e.g. the hydrogen maleate salt.

The pharmaceutical compositions of the present invention are useful in the known indications of the particular active agent incorporated therein.

The exact amounts of active agent and of the formulation to be administered depend on a number of factors, e.g. the condition to be treated, the desired duration of treatment and the rate of release of active agent.

For example, the amount of the active agent required and the release rate thereof may be determined on the basis of conventional in vitro or in vivo techniques, determining how long a particular active agent concentration in the blood plasma remains at an acceptable level for a therapeutic effect.

Examples of doses provided in a solid formulation, e.g., a tablet, are, for Irritable Bowel Syndrome (IBS), 1 mg to 12 mg of active agent, for functional dyspepsia (FD) and gastroesophageal reflux disease (GERD), 0.2 to 2mg of active agent, in particular compound A, e.g. the hydrogen maleate salt, per day for a 70 kilogram mammal, e.g. humans, and in standard animal models. The increased tolerability of the active agent, in particular compound A, e.g. the hydrogen maleate salt, provided by the compositions may be observed in standard animal tests and in clinical trials.

The pharmaceutical composition of the invention comprising a 5-HT₄ receptor agonist, partial agonist or antagonist is particularly useful for improving sensory perception of rectal distension, e.g. for the treatment of anal incontinence, or for preventing, modulating or treating visceral pain or discomfort.

5-HT₄ receptor agonists, partial agonists or antagonists, e.g. as disclosed in EP-A1-505,322, on the basis of observed activity, e.g. stimulatory effect on the peristaltic reflex in the isolated guinea-pig ileum, e.g. as described in EP-A1-505,322, have been found to be useful for the treatment of gastro-intestinal motility disorders, for example to normalise or to improve the gastric emptying and intestinal transit in subjects having a disturbed motility, e.g. in irritable bowel syndrome.

In accordance with the present invention, it has now surprisingly been found that 5-HT₄ receptor agonists, partial agonists or antagonists have a beneficial effect, e.g. they exert modulating effects, on the sensory perception of rectal distension and on visceral sensitivity or perception.

It is admitted that receptor properties are not uniform throughout the gut and that the type of afferent innervation reflects the quality of sensations originating from a particular organ. For example, the rectum belongs to those parts of the gastro-intestinal tract from which also non-painful sensations arise, in contrast to the colon from which only painful sensations emanate.

Anal incontinence may be due to functional disturbances of the main anal continence mechanisms. Anal continence appears to be based on a co-ordinated functioning of the

neuromuscular machinery managing rectal sensation and compliance, the recto-anal inhibitory reflex, reflex contractions of the external anal sphincter and the puborectalis muscle. Although skeletal muscle (external sphincter and puborectalis) contractions are of great importance in the maintenance of continence, it is probably the triggering effect of rectal sensation and perception that plays a crucial role and, in fact, is frequently abnormal in incontinent patients. Anal incontinence is a dysfunction which occurs particularly in diabetics and the elderly population.

There is a medical need for modulating visceral sensitivity, discomfort or pain in patients suffering from gastro-intestinal disorders and for a treatment of anal continence dysfunctions.

In accordance with the particular findings of the present invention, there is provided:

- 1.1. A method for preventing, modulating or treating visceral, e.g. abdominal, pain or discomfort in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof.
- 1.2. A method for modulating visceral sensitivity or perception in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof.
- 1.3. A method for stimulating 5-HT₄ receptors present on afferent nerve terminals, particularly on extrinsic neurones of the gut, in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄ receptor agonist or partial agonist or a pharmaceutically acceptable salt thereof.
- 1.4. A method for modulating visceral sensitivity, discomfort or pain via stimulation of 5-HT₄ receptors present on afferent nerve terminals, particularly on extrinsic neurones of the gut, in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄ receptor agonist or partial agonist or a pharmaceutically acceptable salt thereof.

- 1.5. A method for regulating or stabilising myenteric plexus-afferent fibbers in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄ receptor agonist or partial agonist or a pharmaceutically acceptable salt thereof.
- 1.6. A method for improving sensory perception of rectal distension in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof.
- 1.7. A method for treating anal continence dysfunctions in a subject in need thereof, which method comprises administering to said subject an effective amount of 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof.

As alternative to the above the present invention also provides:

- 2. A 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof for use in a method as defined under 1.1 to 1.7 above; or
- 3. A 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof for use in the manufacture of a pharmaceutical composition for use in a method as defined under 1.1 to 1.7 above; or
- 4. A pharmaceutical composition for use in a method as defined under 1.1 to 1.7 above comprising a 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof, together with one or more pharmaceutically acceptable diluents or carriers therefor, e.g. a composition such as disclosed hereinabove.

Preferred compounds for use in accordance with the invention include e.g. those listed hereinabove, particularly 5-HT₄ receptor full agonists or partial agonists, e.g. (S)-zacopride, cisapride, prucalopride, SB 205149, SC 53116, RS 67333, RS 67506, BIMU 1, BIMU 8, (S)-RS 56532, especially Compound A and particularly its hydrogen maleate salt, more

preferably selective 5-HT₄ receptor agonists or partial agonists, and 5-HT₄ receptor antagonists, e.g. Tropisetron, GR 113808, GR 125487, SB 204070, SB 207266, RS 23597, RS 39604, RS 100235, DAU 6285, SC 53606, 3-(5-hydroxy-7-methyl-1H-indol-3-yl-methylene)-N-pentyl-N-methyl-carbazimidamide etc. By selective is meant a compound which does not substantially bind to or stimulate the serotonin 5-HT₃ receptor. A group of compounds excludes Tropisetron.

Utility of a 5-HT₄ receptor agonist, partial agonist or antagonist in the prevention, modulation or treatment of visceral, *e.g.*, abdominal pain or discomfort or modulation of visceral sensitivity or perception or regulation or stabilisation of myenteric plexus-afferent fibers, is demonstrated in convenient tests, *e.g.*, in accordance with the method hereinafter described.

Decerebrate, anaesthesia-free cats under continuous monitoring of blood pressure are paralysed by alcuronium chloride dissolved in rheomacrodex i.v. (200 μ g/kg initially and supplementary doses of 100 μ g/kg, if necessary), and artificially ventilated. Single unit activity of afferent fibres are recorded in a monopolar fashion from peripheral endings of centrally cut filaments of sacral dorsal roots. Tension receptors are identified by probing of their receptive fields in the wall of the mobilised rectum. Thereafter, the response of the units to barostat -controlled rectal ramp-distension is determined. The quantitative response characteristics of the units is evaluated with respect to distension pressure and resulting rectal diameter. Alternatively, the response of the units to pressure-induced peristalsis is measured.

After obtaining 2 distension profiles (5 min each) and/or 10 min of peristalsis under control conditions, a 5-HT₄ receptor agonist, partial agonist or antagonist, e.g., Compound A, or vehicle is applied i.v. and the protocol is repeated. Subsequently, the activity of additional units is recorded in the presence of a 5-HT₄ receptor agonist, partial agonist or antagonist, e.g., Compound A, or vehicle according to the distension/peristalsis protocol. In this assay, the firing rate of the rectal afferents is reduced after administration of a 5-HT₄ receptor agonist or partial agonist at a dose range of from 0.1 to 3 mg/kg i.v., at distension pressures above 20 mmHg. With Compound A, administered i.v. in incremental doses from 0.15 to 1.2

mg/kg, the most prominent inhibition occurs at 50 mmHg and a half-maximal reduction is obtained at about 0.7 mg/kg.

Utility of a 5-HT₄ receptor agonist, partial agonist or antagonist, e.g., Compound A, in the treatment of anal incontinence as well as utility in treating conditions as hereinabove specified, may be demonstrated in accordance with the method hereinafter described.

Intraluminal pressures and reflexes in the last 60 cm of the colon of 10 fasted healthy volunteers are measured by means of perfusion manometry. Three latex balloons positioned at 50, 30 and 10 cm, allow volume stimulation. Basal values of colonic intraluminal pressures and reflexes are established. Subsequently, reflex inhibitory relaxations of the internal anal sphincter is triggered by inflating the balloons by 10 ml increments up to a maximum volume of 150 ml. During the inflation phase, two parameters are evaluated: a) the reflux threshold (volume able to induce a substantial pressure decrease of the internal anal sphincter); and b) the sensation threshold (volume able to induce a conscious defecation reflex). After the basal recordings, each subject is given a 5-HT₄ receptor agonist, partial agonist or antagonist, e.g., Compound A, p.o. and 30 to 90 min later the colonic intraluminal pressure and reflexes are assessed again by the same method. In this test, the 5-HT₄ receptor agonist, partial agonist or antagonist, e.g., Compound A, significantly reduced the sensation threshold when administered at a dose of 2-12 mg p.o.

5-HT₄ receptor agonists, partial agonists or antagonists, *e.g.*, Compound A, may be administered by any conventional route, in particular enterally, preferably orally, *e.g.*, in the form of tablets or capsules, or parenterally, *e.g.*, in the form of injectable solutions or suspensions or in a suppository form.

5-HT₄ receptor agonists, partial agonists or antagonists, *e.g.*, Compound A, may be administered in free form or in pharmaceutically salt form. Such salts exhibit the same order of activity as the 5-HT₄ receptor agonists, partial agonists or antagonists in free form.

Daily dosages required in practising the method of the present invention will vary depending upon, for example, the particular compound employed, the mode of administration and the

severity of the condition to be treated. An indicated daily dose is in the range of from about 0.05 to about 30 mg, e.g., from about 0.05 to about 5 mg for parenteral use, and of from about 0.1 to about 30 mg for oral use, conveniently administered once or in divided dosages 2 to 4x/day, or in sustained release form. Unit dosage forms for oral administration accordingly comprise from about 0.5 to about 30 mg of 5-HT₄ receptor agonist, partial agonist or antagonist, e.g., Compound A, or a pharmaceutically acceptable salt thereof, admixed with an appropriate solid or liquid, pharmaceutically acceptable diluent or carrier therefor.

Furthermore, it has also been found that a 5-HT₄ receptor agonist or partial agonist-e.g., Compound A, have a beneficial effect in the prevention or treatment of gastro-intestinal motility disorders, e.g. a stimulatory effect on gastrointestinal motility, in horses and cattle.

Accordingly, there is also provided:

- 5.1. A method for preventing or treating gastro-intestinal motility disorders, e.g. by stimulating the motility of the gastro-intestinal tract in horses or cattle in need thereof, which method comprises administering to the horses or cattle an effective amount of a 5-HT₄ receptor agonist or partial agonist, e.g., Compound A, or a pharmaceutically acceptable salt thereof.
- 5.2. A method for preventing or treating gastro-intestinal motility disorders, e.g. after colic surgery, e.g. post-operative lleus, in horses or cattle in need thereof, which method comprises administering to the horses or cattle an effective amount of a 5-HT₄ receptor agonist or partial agonist, e.g., Compound A, or a pharmaceutically acceptable salt thereof.
- 6. A 5-HT₄ receptor agonist or partial agonist, e.g., Compound A, or a pharmaceutically acceptable salt thereof, for use as a veterinary pharmaceutical e.g. for horses or cattle, e.g. in any of the method 5.1 or 5.1 indicated above or for use in the manufacture of a veterinary pharmaceutical e.g. for use in a method as defined under 5.1 or 5.2.

7. A pharmaceutical composition for veterinary use, e.g. in horses or cattle, e.g. in any of the method 5.1. or 5.2. as indicated above, comprising a 5-HT₄ receptor agonist or partial agonist, e.g., Compound A, or a pharmaceutically acceptable salt thereof, together with a pharmaceutically acceptable diluent or carrier therefor, e.g. a composition as disclosed hereinabove.

Preferred 5-HT₄ receptor agonists or partial agonists for use in horses or cattle in accordance with the invention include e.g. those listed hereinabove, e.g. (S)-zacopride, prucalopride, SB 205149, SC 53116, RS 67333, RS 67506, BIMU 1, BIMU 8, (S)-RS 56532, especially Compound A and particularly its hydrogen maleate salt, more preferably a selective 5-HT₄ receptor agonist or partial agonist.

Utility of a 5-HT₄ receptor agonist or partial agonist, *e.g.*, Compound A, in the treatment of post-operative Ileus as well as utility in treating conditions as hereinabove specified in horses or cattle, may be demonstrated in accordance with the method hereinafter described.

20 horses having colic syndrome are submitted to abdominal surgery. During surgery supportive therapy is applied to them. At the end of surgery, a specific 5-HT₄ receptor agonist or partial agonist, *e.g.*, Compound A, is administered i.v. or i.m., e.g. at a dose of from 0.01 to 10 mg/kg. This dose is repeated every 8 to 24 h until spontaneous defecation is observed. Gastro-intestinal motility is evaluated based e.g. on the presence or absence of gastric reflux as determined by nasogastric intubation, occurrence of borborygmi and timing of defecation after the first injection of the test compound. In this test, the compounds tested, *e.g.* Compound A, are effective in restoring normal motility function of the equine intestine.

Daily dosages required in practising the veterinary method of the present invention will vary depending upon, for example, the particular compound employed, the mode of administration and the severity of the condition to be treated. An indicated daily dose is in the range of from about 0.01 to about 10 mg/kg, e.g., from about 0.05 to about 5 mg/kg for parenteral use, conveniently administered once or in divided dosages 2 to 4x/day, or in sustained release form.

In a further aspect the invention provides a method for preventing or treating gastrointestinal motility disorders in a subject, e.g., a human or an animal, in need of such a therapy comprising administering to this subject an effective amount of a composition according to the present invention.

In a further aspect the invention a process is provided for improving dissolution properties in aqueous media of a pharmaceutical composition containing an acid sensitive and/or poorly soluble in aqueous media active agent, e.g., a 5-HT₄ receptor agonist, more particularly compound A, e.g. the hydrogen maleate salt.

The pharmaceutical composition of the invention may be prepared by any conventional method known in the art, e.g., by mixing an appropriate amount of the active agent, e.g., a 5-HT₄ receptor agonist, with at least 15%, e.g., from 20 to 60%, e.g., from 30 to 50%, e.g., 40%, by weight of a disintegrant based on the total weight of the composition.

It is preferred to formulate in solid form, e.g., unit dosage form. Typical form include capsules and preferably compressed forms such as tablets.

The pharmaceutical composition according to the invention may be prepared by e.g., a wet, e.g., water based, granulation manufacturing process (the process equipment, as glass material, may be pre-treated with a siliconizing agent) comprising the successive steps of :

- i) pre-mixing the acid sensitive and/or poorly soluble in water active agent, e.g., a 5-HT₄ receptor agonist, e.g., compound A, e.g. the hydrogen maleate salt with 60 to 98% of the diluent, and then sieving the resulting mixture,
- ii) mixing purified water with the binder in a weight ratio of from 1:20 to 3:20, and stirring until dissolution,
- iii) adding the surfactant to the solution of ii) and stirring until dissolution,
- iv) adding the disintegrant, the remaining diluent and 50 to 70% of the first lubricant to the pre mixture of i) and mixing
- v) wetting the mixture of step iv) with the granulating solution from step iii) while mixing

- vi) granulating the mixture of step v) by mixing,
- vii) drying the granulate to reach a required loss on drying, e.g., for the tabletting mixture viii) sizing the granulate by sieving.

For tablet manufacturing the granulate from viii) is mixed, e.g., in a free fall mixer, with the rest of the first lubricant and the second one to obtain the desired final tabletting mixture which may be compressed into tablets. This may be performed with conventional tabletting machines on, e.g., a rotary machine, at compression pressures of, e.g., 2 to 30 KN, e.g. 5 to 27 KN, e.g., 10 to 20 KN (KN = Kilo Newtons).

The composition according to the invention may also be prepared by an alternative wet granulation manufacturing process wherein the pre-mixing and sieving of step i) are not performed. In this case, the active agent, the disintegrant, the diluent and about 60% of the first lubricant are pre-blended together and then wetted with the wetting solution of step iv).

Compositions comprising any of the above-mentioned active agents may be prepared by a process as disclosed above.

If desired the pharmaceutical compositions of the invention are stored under low relative humidity conditions, e.g., rH (relative humidity) less than 50%, e.g., below e.g. 30-50%, and at room temperature, preferably less than 20°C. The compositions provide storage stable systems. Insignificant degradation is detected after storage of up to 1 year at room temperature, e.g., 25 °C.

The compositions of the invention may be packed in conventional manner to keep out humidity, e.g., in a blister pack, optionally with a desiccant.

The compositions of the invention may have a water content of from 0 to 3% based on the total weight of the composition.

The present invention relates in a further aspect to a composition, in particular comprising compound A, as obtained by one of the above processes to provide a small, stable form.

Examples

The following examples illustrate the manufacturing, on an industrial scale, of compositions comprising compound A hml using a wet granulation process as disclosed above.

Example 1

A 2 mg tablet formulation may be prepared as described hereinafter.

a) Preparation of the granulated material

Premixing step

- 1. 4.432 kg of compound A hml and 28.688 kg of lactose monohydrate are mixed with an intensive mixer (Colette Gral® 300 I or Fielder®); mixer speed setting: 1; chopper speed setting: 1) for approximately 1.5 minutes, or with a free fall mixer (Turbula®, Soneco® or Röhnrad®)
- 2. The pre-mixture from step 1 is then sieved (oscillating granulator, e.g., Frewitt® or Erweka®; mesh size: 0.8 millimetres).
- 3. The pre-mixture is divided into two portions of 16.560 kg.

Preparation of the granulating solution

- 4. Approximately 40 kg of purified water are weighed out.
- 5. 3.600 kg of methylhydroxypropylcellulose 3 maps are added to the purified water from step 4 and this is stirred until dissolution.
- 6. 1.440 kg of poloxamer 188 are added to the solution from step 5 while stirring until dissolution.

Granulating step

- 7. 28.800 kg of crospovidone, 10.080 kg of lactose monohydrate and 4.320 kg of glyceryl monostearate are weighed out.
- 8. One portion of the premixture from step 3 is added to the excipients from step 7 and this is mixed with the intensive mixer, e.g., Colette Gral® 300 I or Fielder® (mixer speed setting: 1; chopper speed setting: 1) for approximately 2 minutes.
- 9. The mixture from step 8 is wetted with the granulating solution from step 6 while mixing with the intensive mixer, e.g., Colette Gral® 300 I or Fielder® (mixer speed

- setting: 1; chopper speed setting: 0; pumping rate approximately: 4 kg/minute) for approximately 12 minutes.
- 10. Approximately 2 kg of purified water are weighed out.
- 11. The vessel from step 6 is rinsed with the purified water from step 10 and this is added to the mixture from step 9 while mixing.
- 12. The mass is granulated by mixing with the intensive mixer, Colette Graf® 300 I or Fielder® (mixer speed setting: 1; chopper speed setting: 1) for approximately 2.5 minutes.

Drying step

- 13. The granulate from step 12 is dried in a fluidised air bed drier (e.g., Glatt[®] or Aeromatic[®]) for approximately 65 minutes (inlet air temperature approximately 70°C) to reach the required loss on drying (LOD) for the tabletting mixture, i.e., until LOD ≤4,4%.
- 14. The granulate sized by sieving (0.8 millimetres) with an oscillating sieve granulator, e.g., Frewitt® or Erweka®.
- 15. Steps 4 to 14 are repeated with the other portion of step 3.

b) Preparation of the tabletting mixture

- 16. 8.640 kg of polyethylene glycol 4000 and 5.760 kg of glyceryl monostearate are sieved (oscillating granulator, e.g., Frewitt® or Erweka®; mesh size: 0.8 millimetres)
- 17. The ingredients from step 16 are added to the total mass of granulated material and this is mixed with a free fall mixer, e.g., Soneco® or Röhnrad®, for approximately 20 minutes (10 rpm) to obtain the desired final tabletting mixture.

c) Compression step

18. The tabletting mixture from step 17 is pressed into tablets using compression pressures of 11, 14 or 17 KN on a rotary tabletting machine, e.g., Fette[®], Korsh[®], Kelian[®] or Coarty[®] (temperature < 20°C; rH (relative humidity) < 40%)</p>

Example 2: Composition of a 2 mg tablet (1 mg of base corresponds to 1.385 mg of the hydrogen maleate salt of compound A)

Compound A hml	2.77 (2mg base)
Polyplasdone XL USP/NF	36.00
Glyceryl monostearate USP/NF	9.00
Poloxalkol	1.80
Lactose 200 mesh	30.53
HPMC 3cPs	4.50
Polyethyleneglycol 4000	5.40
Water adsorbed	2.00
Total	92 mg

Example 3

A 6 mg tablet formulation may be prepared by the manufacturing process described hereinafter.

a) Preparation of the granulated material

Preparation of the granulating solution

- 1. Approximately 40 kg of purified water are weighed out.
- 2. 3.600 kg of methylhydroxypropylcellulose 3 maps are added to the purified water from step 1 while stirring until dissolution.
- 3. 1.440 kg of poloxamer 188 are added to the solution from step 2 while stirring until dissolution (mixing tank under stirring).

Granulating step

- 4. 4.787 kg of compound A hml and 28.800 kg of crospovidone, 21.853 kg of lactose monohydrate and 4.320 kg of glyceryl monostearate are weighed out.
- 5. The ingredients from step 4 are mixed with the intensive mixer, e.g., Colette Gral® 300 I or Fielder® (mixer speed setting: 1; chopper speed setting: 1) for approximately 2 minutes.
- 6. The mixture from step 5 is wetted with the granulating solution from step 3 while mixing with the intensive mixer, e.g., Colette Gral® 300 I or Fielder® (mixer speed

- setting: 1; chopper speed setting: 0; pumping rate approximately 4 kg/minute) for approximately 12 minutes.
- 7. Approximately 2 kg of purified water are weighed out.
- 8. The vessel from step 3 is rinsed with the purified water from step 7 and this is added to the mixture from step 6 while mixing.
- 9. The mass is granulated by mixing with the intensive mixer, e.g., Colette Graf 300 I or Fielder (mixer speed setting: 1; chopper speed setting: 1) for approximately 2.5 minutes.

Drying step

- 10. The granulate from step 9 is dried in a fluidised air bed drier, e.g., Glatt[®] or Aeromatic[®]) for approximately 65 minutes (Inlet air temperature approximately 70°C) to reach the desired loss on drying (LOD) for the tabletting mixture, i.e., until LOD ≤4,4%.
- 11. The granulate sized by sieving (0.8 millimetres) with an oscillating sieve granulator (Frewitt® or Erweka®)
- 12. Steps 1 to 11 are repeated.

b) Preparation of the tabletting mixture

- 13. 8.640 kg of polyethylene glycol 4000 and 5.760 kg of glyceryl monostearate are sieved with an oscillating sieve granulator, e.g., Frewitt® or Erweka® (0.8 millimetres)
- 14. The ingredients from step 13 are added to the total mass of granulated material and this is mixed with a free fall mixer, e.g., Soneco® or Röhnrad®, for approximately 20 minutes (10 rpm) in the desired final tabletting mixture.

c) Compression step

15. The tabletting mixture from step 14 is pressed into tablets using compression pressures of 13, 16 or 19 KN on a rotary tabletting machine, e.g., Fette[®], Korsh[®], Kelian[®] or Coarty[®] (temperature<20°C, rH (relative humidity) < 40 %).

Example 4: Composition of a 6 mg tablet (1 mg of base corresponds to 1.385 mg of hydrogen maleate of compound A):

Compound A hml	8.31 (6mg base)
Polyplasdone XL USP/NF	50.00
Glyceryl monostearate USP/NF	12.50
Poloxalkol	2.50
Lactose 200 mesh	37.94
HPMC 3cPs	6.25
Polyethyleneglycol 4000	7.50
Water adsorbed	3.00
Total	128 mg

Example 5

A 0.5 mg tablet formulation may be prepared by the manufacturing process described hereinafter.

a) Preparation of the granulated material

Premixing step

- 1.994 kg of compound A hml and 31.126 kg of lactose monohydrate are mixed with an intensive mixer (Colette Gral® 300 I or Fielder®); mixer speed setting: 1; chopper speed setting: 1) for approximately 1.5 minutes, or with a free fall mixer (Turbula®, Soneco® or Röhnrad®)
- 2. The premixture from step 1 is then sieved (oscillating granulator, e.g., Frewitt® or Erweka®; mesh size: 0.8 millimetres).
- 3. The premixture is divided into two portions of 16.560 kg.

Preparation of the granulating solution

- 4. Approximately 43 kg of purified water are weighed out.
- 5. 3.600 kg of methylhydroxypropylcellulose 3 maps are added to the purified water from step 4 and this is stirred until dissolution.
- 6. 1.440 kg of poloxamer 188 are added to the solution from step 5 while stirring until dissolution.

Granulating step

- 7. 28.800 kg of crospovidone, 10.080 kg of lactose monohydrate and 4.320 kg of glyceryl monostearate are weighed out.
- 8. One portion of the premixture from step 3 is added to the excipients from step 7 and this is mixed with the intensive mixer, e.g., Colette Gral® 300 I or Fielder® (mixer speed setting: 1; chopper speed setting: 1) for approximately 2 minutes.
- 9. The mixture from step 8 is wetted with the granulating solution from step 6 while mixing with the intensive mixer, e.g., Colette Gral® 300 I or Fielder® (mixer speed setting: 1; chopper speed setting: 0; pumping rate approximately: 4 kg/minute) for approximately 12 minutes.
- 10. Approximately 2 kg of purified water are weighed out.
- 11. The vessel from step 6 is rinsed with the purified water from step 10 and this is added to the mixture from step 9 while mixing.
- 12. The mass is granulated by mixing with the intensive mixer, Colette Graf® 300 I or Fielder® (mixer speed setting: 1; chopper speed setting: 1) for approximately 2.5 minutes.

Drying step

- 13. The granulate from step 12 is dried in a fluidised air bed drier (e.g., Glatt® or Aeromatic®) for approximately 60 minutes (inlet air temperature approximately 70°C) to reach the required loss on drying (LOD) for the tabletting mixture, i.e. until LOD ≤4,5%.
- 14. The granulate sized by sieving (0.8 millimetres) with an oscillating sieve granulator, e.g., Frewitt® or Erweka®.
- 15. Steps 4 to 14 are repeated with the other portion of step 3.

b) Preparation of the tabletting mixture

- 16. 8.640 kg of polyethylene glycol 4000 and 5.760 kg of glyceryl monostearate are sieved (oscillating granulator, e.g., Frewitt® or Erweka®; mesh size: 0.8 millimetres)
- 17. The ingredients from step 16 are added to the total mass of granulated material and this is mixed with a free fall mixer, e.g., Soneco® or Röhnrad®, for approximately 20 minutes (10 rpm) to obtain the desired final tabletting mixture.

c) Compression step

Total

18. The tabletting mixture from step 17 is pressed into tablets on a rotary tabletting machine, e.g., Fette[®], Korsh[®], Kelian[®] or Coarty[®] (temperature < 20°C; rH (relative humidity) < 40%)

Example 6: Composition of a 0.5 mg tablet (1 mg of base corresponds to 1.385 mg of the hydrogen maleate salt of compound A)

Compound A hml	0.6925 (0.5mg base)
Polyplasdone XL USP/NF	20.00
Glyceryl monostearate USP/NF	5.00
Poloxalkoi	1.00
Lactose 200 mesh	17.8075
HPMC 3cPs	2.50
Polyethyleneglycol 4000	3.00
Water adsorbed	1.00

Example 7: Composition of a 12 mg tablet (1 mg of base corresponds to 1.385 mg of the hydrogen maleate salt of compound A)

51 mg

The manufacturing process is similar to the process used for the 6mg tablets.

Compound A hml	16.62 (12mg base)
Polyplasdone XL USP/NF	72.00
Glyceryl monostearate USP/NF	18.00
Poloxalkol	3.60
Lactose 200 mesh	49.98
HPMC 3cPs	9.0
Polyethyleneglycol 4000	10.8
Water adsorbed	4.00
Total	184 mg

Claims:

- A solid oral pharmaceutical composition comprising an effective amount of an acid sensitive active agent and a disintegrant which is present in an amount of at least 15% by weight based on the total weight of the composition.
- 2. A solid oral pharmaceutical composition comprising an effective amount of an active agent which is poorly soluble in aqueous media and a disintegrant which is present in an amount of at least 15% by weight based on the total weight of the composition.
- 3. A pharmaceutical composition according to claim 1 or 2 wherein the active agent has a solubility in aqueous media less than 1%.
- 4. A pharmaceutical composition according to any of claims 1 to 3 wherein the active agent is a serotonergic compound.
- 5. A pharmaceutical composition as claimed in any preceding claim wherein the active agent is a 5-HT₄ receptor antagonist.
- 6. A pharmaceutical composition as claimed in any of claims 1 to 4 wherein the active agent is a 5-HT₄ receptor agonist.
- 7. A pharmaceutical composition according to claim 6 wherein the 5-HT₄ receptor agonist is Tegaserod, preferably its hydrogen maleate (hml) salt.
- 8. A pharmaceutical composition as claimed in any preceding claim wherein the disintegrant is crospovidone.
- A pharmaceutical composition as claimed in any preceding claim comprising a lubricant.
- A pharmaceutical composition according to claim 9 wherein the lubricant comprises a glyceryl mono fatty acid.

- 11. A pharmaceutical composition according to claim 9 wherein the lubricant comprises a mixture of glyceryl monostearate and polyethylene glycol.
- A pharmaceutical composition as claimed in any preceding claim comprising a surfactant.
- 13. A pharmaceutical composition according to claim 12 wherein the surfactant comprises poloxamer.
- 14. Use of at least 15% by weight of a disintegrant in the manufacturing of a solid pharmaceutical composition for the administering of an acid sensitive active agent.
- 15. Use of at least 15% by weight of a disintegrant in the manufacturing of a solid pharmaceutical composition for the administering of an active agent being acid sensitive and/or having a poor water solubility.
- 16. Use according to claim 14 or 15 wherein the active agent is a 5-HT₄ receptor agonist.
- 17. Use according to claim 16 wherein the 5-HT₄ receptor agonist is Tegaserod, preferably its hydrogen maleate salt.
- 18. Use of a pharmaceutical composition according to any one of claims 1 to 13 for the the manufacture of a composition for the prevention and treatment of gastro-intestinal motility disorders in humans or animals.
- 19. A process for improving dissolution properties of a pharmaceutical composition as claimed in any of claims 1 to 13.
- 20. A method for preventing, modulating or treating visceral pain or discomfort, for modulating visceral sensitivity or perception, for improving sensory perception of rectal distension, or for treating anal continence dysfunctions in a subject in need thereof, which method comprises administering to said subject an effective amount of a 5-HT₄

receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof.

- 21. A 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof for use in the manufacture of a pharmaceutical composition for use in preventing, modulating or treating visceral pain or discomfort, modulating visceral sensitivity or perception, improving sensory perception of rectal distension, or treating anal continence dysfunctions.
- 22. A pharmaceutical composition for use in preventing, modulating or treating visceral pain or discomfort, modulating visceral sensitivity or perception, improving sensory perception of rectal distension, or treating anal continence dysfunctions, which composition comprises a 5-HT₄ receptor agonist, partial agonist or antagonist or a pharmaceutically acceptable salt thereof, together with one or more pharmaceutically acceptable diluents or carriers therefor.
- 23. A method for preventing or treating gastro-intestinal motility disorders in horses or cattle in need thereof, which method comprises administering to the horses or cattle an effective amount of a 5-HT₄ receptor agonist or partial agonist or a pharmaceutically acceptable salt thereof.
- 24. A 5-HT₄ receptor agonist or partial agonist or a pharmaceutically acceptable salt thereof, for use as a veterinary pharmaceutical or for use in the manufacture of a veterinary pharmaceutical.
- 25. A pharmaceutical composition for veterinary use comprising a 5-HT₄ receptor agonist or partial agonist or a pharmaceutically acceptable salt thereof, together with a pharmaceutically acceptable diluent or carrier therefor.
- 26. A pharmaceutical composition comprising Tegaserod having dissolution characteristics in water or USP buffers pH 6.8 and 7.5 of :

time (minutes)	amount (percentage)
5	30 - 90
15	80 - 100
30	95 - 100
60	100

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7: A61K 9/20, A61P 1/00		(11) International Publication Number	er: WO 00/10526	
A61K 9/20, A61F 1/00	A3	(43) International Publication Date:	2 March 2000 (02.03.00)	
(21) International Application Number: PC	г/EP99/060	Pommard, F-75012 Paris (FI Heuwinkelstrasse 13, CH-41	R). ZÜGER, Othmar [CH/CH]; 23 Allschwil (CH).	
(22) International Filing Date: 19 August 19	99 (19.08.9			

(30)) Priority	Data:
(JU	,	vau.

9818340.3 9823477.6	21 August 1998 (21.08.98) 27 October 1998 (27.10.98)	GB GB
9910320.2	5 May 1999 (05.05.99)	GB
9911059.5	12 May 1999 (12.05.99)	GB

- (71) Applicant (for all designated States except AT US): NOVAR-TIS AG [CH/CH]; Schwarzwaldallee 215, CH-4058 Basel (CH).
- (71) Applicant (for AT only): NOVARTIS-ERFINDUNGEN VER-WALTUNGSGESELLSCHAFT MBH [AT/AT]; Brunner Strasse 59, A-1230 Vienna (AT).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): DE BRUIJN, Karel [NL/FR]; 13, rue du Muhlberg, F-68730 Blotzheim (FR). ENGEL, Günter [DE/DE]; Im Hasengarten 11, D-79576 Weil (DE). PFANNKUCHE, Hans-Jürgen [DE/DE]; Vierthauen 17, D-79576 Weil (DE). THEWIS-SEN, Michael [DE/DE]; Apollinarisstrasse 26, D-40227 Düsseldorf (DE). VITZLING, Christian [FR/FR]; 39, rue de

- (74) Agent: BECKER, Konrad; Novartis AG, Corporate Intellectual Property, Patent & Trademark Dept., CH-4002 Basel (CH).
- (81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(88) Date of publication of the international search report: 8 September 2000 (08.09.00)

(54) Title: NEW ORAL FORMULATION FOR 5-HT4 AGONISTS OR ANTAGONISTS

(57) Abstract

The present invention relates to a pharmaceutical composition, in particular to a composition for administering active agents which are poorly soluble in aqueous media, and/or which are acid sensitive.

> U.S. Serial No.: 09/900,336 Docket No.: 484482000300

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑÜ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Yugoslavia Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	211	Zimoabwe
СМ	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	Li	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG			
			2.00112	30	Singapore		

Intel onal Application No PCT/EP 99/06083

A. CLASSIFICATION F SUBJECT MATTER IPC 7 A61K9/20 A61P A61P1/00 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) IPC 7 A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ° Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X CHEMICAL ABSTRACTS, vol. 127, no. 15, 1-4,6,9,13 October 1997 (1997-10-13) 12, Columbus, Ohio, US; 14-16. abstract no. 210377, 18,19 OUCHI, HIROSHI ET AL: "Digestive tract disease-treating agents" XP002129585 abstract & JP 09 194374 A (TAISHO PHARMACEUTICAL CO., LTD., JAPAN) 29 July 1997 (1997-07-29) X US 5 523 289 A (ALVAREZ FRANCISCO J ET 1,8,9, AL) 4 June 1996 (1996-06-04) 12,14 column 23 -column 24; example 12 X Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 24 May 2000 14 06, 2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Boulois, D Fax: (+31-70) 340-3016

Inter anal Application No PCT/EP 99/06083

C.(Continu	uation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 96 16639 A (CIBUS PHARMACEUTICAL INC) 6 June 1996 (1996-06-06) page 32 -page 33	2,3,8,9, 12,15
X	WO 97 22335 A (ABBOTT LAB) 26 June 1997 (1997-06-26) page 8; table 1	1-3,9, 12,14,15
X	PATENT ABSTRACTS OF JAPAN vol. 013, no. 419 (C-637), 18 September 1989 (1989-09-18) & JP 01 156909 A (CHUGAI PHARMACEUT CO LTD), 20 June 1989 (1989-06-20) abstract	1-4,14, 15
(EP 0 732 333 A (LILLY CO ELI) 18 September 1996 (1996-09-18) page 19; example 3	1-6,9, 12,15, 16, 18-22, 24,25
K	EP 0 691 340 A (SANWA KAGAKU KENKYUSHO CO) 10 January 1996 (1996-01-10) page 2, line 3 - line 5 page 9, line 10 - line 45 page 10; example 1	1-4,6,9, 12,14-16
(WO 98 03173 A (SEPRACOR INC) 29 January 1998 (1998-01-29) page 5, line 1 - line 8 page 31; example 7	1-4,6,9, 12,14-16
1	AMER. PHARM. ASSOC. / PHARM. SOC. OF GREAT BRITAIN: "Handbook of pharmaceutical excipients" 1988 , AM. PHARM. ASSOC. , WASHINGTON XP002129584 153140 page 289 -page 293	1,2
(EP 0 505 322 A (SANDOZ AG ;SANDOZ LTD (CH); SANDOZ AG (DE)) 23 September 1992 (1992-09-23) cited in the application page 23, line 9 - line 22	20-22, 24,25
(US 5 399 562 A (BECKER DANIEL P ET AL) 21 March 1995 (1995-03-21) column 1, line 6 - line 27	20-22, 24,25
I	FR 2 753 196 A (SYNTHELABO) 13 March 1998 (1998-03-13) page 24, line 21 - line 35	20-22, 24,25
	-/	

Inte onal Application No PCT/EP 99/06083

Category *	citation of decrement, with indication where environment, at the relevant		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
X	STEINER A. ET AL: "Drugs coordinating and restoring gastrointestinal motility and their effects on selected hypodynamic gastrointestinal disorders to horses and cattle" J. OF VET. MED. SERIES A, vol. 42, 1995, pages 613-631, XP000910391 page 617	21,22,24	
X	BRIKAS P.: "Motor modifying properties of 5-HT3 and 5-HT4 receptor agonists on ovine abomasum " J. OF VET. MEDICINE SERIES A, vol. 41, 1994, pages 150-158, XP000910373 page 156 -page 157	20-25	
X	GASTER L.: "SB207266 : 5-HT4 receptor antagonist agent for irritable bowel syndrome" DRUGS FUTURE, vol. 22, no. 12, 1997, pages 1325-1332, XP002099385 page 1328, last paragraph -page 1329	20-22, 24,25	
P,X	LANGAKER KJ ET AL: "The partial 5-HT4 (HTF 919) improves symptoms in constipation predominant irritable bowel syndrome (C-IBS)" DIGESTION, vol. 59, no. 3, 6 September 1998 (1998-09-06), page gapp00064 XP000909235 abstract	20 - 22, 24,25	
X	FARTHING M.: "New drugs in the management of the irritable bowel syndrome" DRUGS, vol. 56, no. 1, July 1998 (1998-07), pages 11-21, XP000909287 page 13; table 2 page 18, paragraph 2	20-25	
A	BERKOW R. ET AL: "The Merck Manual of Diagnosis and Therapy" 1992 , MERCK RESEARCH LABORATORIES , USA XP002138260 215310 page 841 -page 842	20-25	

5

Inter anal Application No PCT/EP 99/06083

		PC1/EP 99/06083				
	(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT					
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.				
A	SCHUTZE K. ET AL: "Double blind study of the effect of cisapride on constipation and abdominal discomfort as component of the irritable bowel syndrome" ALIMENT. PHARMACOL. THER., vol. 11, no. 2, 1997, pages 387-394, XP000909210 page 392 -page 393	20-25				
X	TALLEY N J: "REVIEW ARTICLE: 5-HYDROXYTRYPTAMINE AGONISTS AND ANTAGONISTS IN THE MODULATION OF GASTROINTESTINAL MOTILITY AND SENSATION: CLINICAL IMPLICATIONS" ALIMENTARY PHARMACOLOGY & THERAPEUTICS, GB, BLACKWELL SCIENTIFIC PUBLICATIONS LTD., CAMBRIDGE, vol. 6, no. 3, 1 June 1992 (1992-06-01), pages 273-289, XP000566133 ISSN: 0269-2813 page 277; table 2 page 280 -page 281; tables 3,4	20-22, 24,25				
T	SCOTT & PERRY: "TEGASEROD" DRUGS,AT,ADIS INTERNATIONAL LTD, vol. 58, no. 3, 1999, pages 492-496, XP000874675 ISSN: 0012-6667 the whole document	20-25				
T	GRAUL A ET AL: "TEGASEROD MALEATE" DRUGS OF THE FUTURE,ES,BARCELONA, vol. 24, no. 1, 1999, pages 38-44, XP000874672 ISSN: 0377-8282 the whole document	20-25				

PCT/EP 99/06083

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 1,2,4,5,6,14,15,16,17,26 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: see FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
A. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. X No protest accompanied the payment of additional search fees.

International Application No. PCT/EP 99 \(06083 \)

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box 1.2

Claims Nos.: 1,2,4,5,6,14,15,16,17,26

Present claims 1,2,4,5,6,14,15,16,17, 20-25 relate to compounds defined by their chemical properties, namely "an acid sensitive agent", a agent "poorly soluble in aqueous media", or by their pharmacological profile, namely 5-HT4 agonists or antagonists.

The definition of an acid sensitive agent given on page 2, lines 1-5 and of the poor water solubility of the active on page 1 last paragraph, is not sufficient and too unclear to determine which active agents are concerned, and emcompasses potentially an huge number of compounds, which makes it difficult or impossible to search those compounds. Moreover, a compound cannot be sufficiently characterised by its pharmacological profile or its mode of action as it is done by an expression like serotonergic compound, 5-HT4 agonist or 5-HT-4 antagonist, this expression comprising a lot of compounds having a possible known or unknown action on this receptor.

The claims cover all compounds having this characteristic or property, whereas the application provides support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT for only a very limited number of such compounds. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Independent of the above reasoning, the claims also lack clarity (Article 6 PCT). This lack of clarity in the present case is such as to render a meaningful search over the whole of the claimed scope impossible. Consequently, the search has been carried out for those parts of the claims which appear to be clear, supported and disclosed, namely those parts relating to the compounds described in the present application (see pages 2-4) and disclosed in the examples, and on the concepts "acid sensitive agents" and "poorly soluble agents", "serotonergic" compounds and "5-HT4" agonists or antagonists per se.

Present claim 26 relate to an extremely large number of possible compositions. Support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT is to be found, however, for only a very small proportion of the compositions, namely compositions as defined in claims 1 or 2 and comprising more than 15% of a disintegrant. It is clear that the scope of claim 26 encompasses compositions presenting the same profile, but which are not disclosed in the present application.

In the present case, claim 26 so lacks support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Consequently, the search has been carried out for those parts of the claims which appear to be supported and disclosed, namely those parts relating to the composition comprising tegaserod and a disintegrating agent in a concentration higher than 15% by weight.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant

International Apolication No. PCT/EP 99 06083

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210
is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.
•

International Application No. PCT/EP 99 06083

FURTHER INFORMATION CONTINUED FROM PCT/ISA 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-19,26

Solid oral composition comprising an acid sensitive agent or an agent poorly soluble in aqueous media and at least 15% by weight of the composition of a disintegrant $\frac{1}{2}$

2. Claims: 20-25

A 5-HT4 agonist, antagonist or partial agonist and its use in gastrointestinal disorders

information on patent family members

Inte anal Application No PCT/EP 99/06083

Patent document cited in search repo	n	Publication date		Patent family member(s)	Publication date
JP 9194374	Α	29-07-1997	NON	E	1
US 5523289	A	04-06-1996	AU	2596592 A	21-05-1993
			PT	100976 A	28-02-1994
			WO	9307886 A	29-04-1993
			AT	123282 T	15-06-1995
			AU	646859 B	10-03-1994
			AU	7607791 A	14-11-1991
			CA	2041825 A	12-11-1991
			DE	69110086 D	06-07-1995
			DE	69110086 T	14-12-1995
			DK	456185 T	30-10-1995
4			EP	0456185 A	13-11-1991
			ES	2075261 T	01-10-1995
			GR	3017168 T	30-11-1995
			IE	68045 B	15-05-1996
			IL	97994 A	16-10-1996
			JP	1980629 C	17-10-1995
			JP	4279572 A	05-10-1992
			JP	7000607 B	11-01-1995
			JP	5271194 A	19-10-1993
			MX	9202877 A	30-06-1992
			PT	97631 A,B	31-03-1992
			US	5275950 A	04-01-1994
			US	5344990 A	06-09-1994
			US	5178877 A	12-01-1993
			US US	5310740 A	10-05-1994
				5284849 A 	08-02-1994
WO 9616639	Α	06-06-1996	AU	690417 B	23-04-1998
			AU	4595196 A	19-06-1996
			CA	2205442 A	06-06-1996
			EP FI	0804169 A	05-11-1997
			JP	972305 A	30-05-1997
			NO	10509982 T 972485 A	29-09-1998
			US	5993860 A	31-07-1997 30-11-1999
					30-11-1999
WO 9722335	Α	26-06-1997	US	5705190 A	06-01-1998
			AT	170744 T	15-09-1998
			AU	701268 B	21-01-1999
			UA	1025297 A	14-07-1997
			CA CZ	2209714 A	26-06-1997
			DE	9702212 A 69600620 D	17-12-1997
			DE	69600620 T	15-10-1998
			EP	0799028 A	06-05-1999 08-10-1997
			ES	2122810 T	16-12-1998
			HU	9800516 A	28-08-1998
			JP	11513406 T	28-08-1998 16-11-1999
			NZ	323332 A	27-04-1998
			PL.	321363 A	08-12-1997
			WO	9856357 A	17-12-1998
JP 01156909	Α	20-06-1989	JP	2708803 B	04-02-1998
ED 072222	Α	18-09-1996	US	5654320 A	05-08-1997
EP 0732333		10 07 1330	เมอ		

information on patent family members

Inte. onal Application No PCT/EP 99/06083

Patent document cited in search report		Publication date		atent family member(s)	Publication date
EP 0732333	A	·	AU BR CA CN HU JP NO	6894896 A 9608223 A 2215359 A 1183041 A 9801386 A 11502230 T 974221 A	18-11-1996 29-12-1998 31-10-1996 27-05-1998 28-06-1999 23-02-1999 27-10-1997
·			NZ PL SK WO US US	315971 A 322362 A 124297 A 9633713 A 5798367 A 5817676 A	28-01-1999 19-01-1998 04-02-1998 31-10-1996 25-08-1998 06-10-1998
EP 0691340	A	10-01-1996	JP US	8020586 A 5539125 A	23-01-1996 23-07-1996
WO 9803173		29-01-1998	US AU BG BR CA CN CZ EP JP NO SK US	5739151 A 3592097 A 102757 A 9708287 A 2245768 A 1215992 A 9802508 A 0896539 A 11510828 T 983636 A 123998 A 5877188 A	14-04-1998 10-02-1998 31-08-1999 18-01-2000 29-01-1998 05-05-1999 14-04-1999 17-02-1999 21-09-1999 19-10-1998 10-03-1999 02-03-1999
EP 0505322	A	23-09-1992	AT AU CA CS DE DE ES FI HU IL JP MNO NZ RO SG RU SC VZ	170838 T 651442 B 1309292 A 2063671 A 9200858 A 69226894 D 69226894 T 2121836 T 921222 A 971545 A 64023 A 9500315 A 101312 A 2593022 B 5086026 A 9201244 A 179171 B 242069 A 109194 A 43221 A 279214 B 2095347 C 5510353 A 9202071 A	15-09-1998 21-07-1994 24-09-1992 23-09-1992 14-10-1992 15-10-1998 18-02-1999 16-12-1998 23-09-1992 11-04-1997 29-11-1993 30-10-1995 18-03-1997 19-03-1997 06-04-1993 01-01-1993 13-05-1996 24-02-1995 30-12-1994 17-10-1997 05-08-1998 10-11-1997 23-04-1996 20-09-1993
US 5399562	Α	21-03-1995	NONE		
FR 2753196	Α	13-03-1998	AU	4211997 A	02-04-1998

Information on patent family members

Inte. onal Application No PCT/EP 99/06083

PCT/EP 99/06083 Patent document cited in search report Publication date Patent family member(s) Publication date FR 2753196 Α WO 9811112 A 19-03-1998